

tension sensitive than other myosins with similar length lever arms, suggesting that different myosins have different tension-sensitive transitions.

2896-Pos

Myosin-I Dependent Membrane-Cytoskeleton Adhesion as a Regulator of Cell Surface Morphology

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All cellular functions involving deformation of the plasma membrane (e.g. endocytosis, exocytosis, the formation of surface protrusions) are regulated by the apparent membrane tension (T_m), a composite of the surface tension (σ) and the adhesion provided by molecules linking membrane to the actin cytoskeleton (γ). Using an optical trap based tether force assay, we recently demonstrated that class I myosins, a family of membrane-binding actin-based motor proteins, control membrane tension by mediating membrane-cytoskeleton adhesion. More specifically, these studies revealed that the membrane-cytoskeleton adhesion provided by myosin-I increases apparent membrane tension as indicated by the force required to pull a single membrane tether from the cell surface. Interestingly, the physical links to the cytoskeleton provided by myosin-I also allow the cell to form multiple adjacent tethers. However, when multiple tethers are pulled, tether lifetimes appear to decrease as apparent membrane tension increases. Here we present a thermodynamic model, which accounts for the impact of apparent membrane tension on the lifetime of multiple tethers. We argue that, in the context of our experimental geometry, the global increase in apparent membrane tension that results from greater membrane-cytoskeleton adhesion, works locally to accelerate the rupture of bonds between the membrane and cytoskeleton, which would otherwise prevent individual membrane tethers from coalescing. We also elaborate on this concept to develop a separate model, which shows that tuning the level of membrane-cytoskeleton adhesion may enable cells to vary the density of surface protrusions (i.e. # of structures per unit area membrane). As such, myosin-I dependent membrane-cytoskeleton adhesion emerges as the key regulator of cell surface morphology.

2897-Pos

Kinetics of Myosin-I-Membrane Detachment Under Load

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Myo1c is a single-headed motor that links cell membranes to the underlying actin cytoskeleton. Actin binding occurs via the motor domain, while the tail domain interacts with phosphatidylinositol 4,5-bisphosphate (PI(4,5)P₂) through a putative pleckstrin homology (PH) domain. In this study, we measured the strength and lifetime of the myo1c attachment to spherical supported lipid bilayers (SSL) composed of 1 μ m diameter silica or polystyrene beads coated with various mole fractions of PI(4,5)P₂, dioleoyl-phosphatidylcholine (DOPC), and dioleoyl-phosphatidylserine (DOPS). The SSLs trapped by a laser beam were brought into contact with immobilized spherical (2 μ m) silica pedestals sparsely labeled with the myo1c tail domain. Upon repeated contact and retraction cycles, binding events and subsequent rupture forces were measured. The most probable rupture forces (< 10 pN) from membranes containing 2% PI(4,5)P₂ were found to be largely independent of the loading rate (pN/s). Nevertheless, the frequency of single myo1c molecules interacting with membranes containing 2% PI(4,5)P₂ - 98% DOPC was considerably higher than with membranes that contained 20 - 80% DOPS. Lifetime measurements of myo1c attachment to 2% PI(4,5)P₂ - 98% DOPC membranes under constant pulling forces (1 - 3 pN) were fitted to Bell's equation and the extrapolated duration of the bond at zero force was found to be ~ 25 ms. Although phosphoinositide binding is crucial for the proper cellular targeting of myo1c, our results suggest that it is unlikely that this connection is a suitable anchor for force generation.

2898-Pos

Kinetics and Thermodynamics of Nucleotide Binding Pocket Opening/closing in Myosin V Monitored with FRET

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Kinetic and structural studies of both muscle and non-muscle myosins have revealed that the enzymatic cycle of these motors frequently contains more than one actomyosin ADP state. Interestingly, the rate of ADP release in myosin motors is thought to be the main determinant of sliding velocity in muscle, suggesting strain dependent ADP release may be a critical mechanism of mechanochemical coupling. Our previous work has demonstrated that labeling myosin V in the upper 50 kDa domain with the biarsenical dye FIAsh (MV FIAsh) can serve as an acceptor for fluorescence resonance energy transfer studies with mant labeled nucleotides. We also determined that this donor-acceptor pair likely monitors opening/closing of the nucleotide binding pocket. Currently, we utilized the FRET signal to examine the kinetics of nucleotide binding pocket opening during the process of mantADP release from acto-

MV FIAsh. We obtained evidence that the nucleotide binding pocket goes from a closed to an open conformation prior to the release of ADP. We also explored the temperature dependence of the closed to open transition and nucleotide release steps. We find that at lower temperatures the closed conformation is favored while at higher temperature the open conformation is favored. The more rapid ADP release step which follows nucleotide binding pocket opening is also temperature dependent. Therefore, since both steps are temperature-dependent they likely require significant conformational changes. We also compared our FRET results to the rate of ATP-induced dissociation from actin in the presence of ADP monitored by light scatter. Understanding how strain alters either of these two steps may be critical for elucidating the structural mechanism of strain-dependent ADP release in myosins.

2899-Pos

Coupling the Actin Binding Cleft and Nucleotide Binding Pocket in Myosin V

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Previously we have demonstrated in fluorescence resonance energy transfer (FRET) studies that mant labeled nucleotides and IAEDANS actin can act as good donor probes for a FIAsh labeled acceptor site in the upper 50 kDa domain of myosin V. We examined the temperature dependence of the FRET signal between mantADP and MV WT FIAsh in the presence and absence of actin. We found that at low temperature (4-15°C) a high FRET state dominates (closed pocket) while at high temperature (30-35°C) a low FRET state dominates (open pocket). This transition is reversible suggesting a temperature-dependent conformational change. However, the mutant E442A, which is incapable of hydrolyzing ATP, remains in a high FRET state (closed pocket) with mantATP bound in the presence or absence of actin. Our results suggest a more flexible conformation of myosin in the presence of ADP compared to ATP which allows myosin to populate two actomyosin.ADP state conformations. These results are supported by the lifetime FRET analysis, and by computational FIRST/FRODA analysis of the intrinsic flexibility found in different x-ray crystal structures. We also plan to explore the temperature dependent conformational dynamics of the actin binding cleft using the IAEDANS actin (donor) and MV FIAsh (acceptor) pair in the presence of ATP, ADP, and absence of nucleotide using steady state and lifetime based FRET measurements. Our results will provide critical insights into the mechanocoupling that may occur between the nucleotide-binding pocket and actin binding cleft in myosin motors.

2900-Pos

Interaction of a Class V Myosin from Budding Yeast with its Adapter Protein

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Like their mammalian counterparts, class V myosins in *S. cerevisiae* (Myo2p and Myo4p) bind to various adapter proteins to target a particular cargo for transport. Myo4p uses the adapter proteins She3p and She2p in order to transport mRNA from the mother cell to the bud. She3p binds to the rod of Myo4p, and prevents it from dimerizing, thus forming a single-headed motor complex (Hodges et al., 2008; Bookwalter et al., 2009). Because the Myo4p/She3p complex is single-headed, the question arises as to whether enough motors can bind to a single She2p to enable continuous cargo transport. The She2p crystal structure suggested that She2p exists as a dimer (Niessing et al., 2004). In contrast, our sedimentation equilibrium measurements of She2p were consistent with formation of a tetramer in solution, in principle allowing for binding of four motor heads. We showed that Myo4p/She3p forms a complex with tetrameric She2p in the absence of mRNA, based on sedimentation velocity experiments and co-purification. Mutation of Ser 120 to Tyr converts She2p to a dimer. The ability of the motor complex to bind to this and other She2p mutants is being tested in order to map the binding interface. Total internal reflection fluorescence microscopy is being used to test whether the native She2p tetramer can bind enough single-headed motors to support continuous movement on actin. The ability of She2p mutants to support correct bud tip localization of *ASH1* mRNA in living yeast cells will also be assessed. These studies will help elucidate how a non-processive single-headed motor can act as a cargo transporter.

2901-Pos

The Mechanical Properties of a Single Myosin V Motor Domain During Gait Motion

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